201-15018B

Appendix B – Robust Summaries

Melting Point	B-2
Boiling Point	B-3
Vapor Pressure	B-4
Partition Coefficient	B-6
Water Solubility	B-7
Photodegradation	B-8
Stability in Water	B-10
Transport & Distribution (Fugacity)	B-11
Biodegradation	B-13
Acute Toxicity - Fish	B-15
Acute Toxicity - Invertebrate	B-18
Acute Toxicity – Aquatic Plant	B-19
Acute Toxicity - Mammalian	B-21
Genetic Toxicity (in vitro)	B-31
Genetic Toxicity (in vivo)	B-39
Repeated Dose Toxicity	B-41
Reproductive Toxicity	B-51

Ot JAN -9 PM 1:56

Melting Point

ID: 1

Test Substance Identity: 2-Vinylpyridine
Test Substance Purity: not applicable
Test Substance Remarks: not applicable

Method / Guideline Followed: Modeled using EPI Suite™ v3.11, MPBPWIN v1.41.

GLP: no **Year**: 2003

Test Conditions Remarks: not applicable

Melting Point in °C: -15.16°C (NOTE: < 0°C)

Decomposition: not applicable

Sublimation: not applicable

Results Remarks: Modeled melting point using MPBPWIN v1.41 resulted in a mean melting

point of -15.16°C (mean of Adapted Joback Method and Gold and Ogle Method). This estimate is supported by the fact that Reilly Industries, a supplier of 2-Vinylpyridine, recommends storage of the material below -5°C to maintain product quality; it remains a liquid at these temperatures. (See

Reference #2.)

Conclusions: According to the EPA guidelines, if the estimated melting point is below

0°C, there is no need to provide measured data for this endpoint. (See

Reference #3.)

Conclusions Remarks: not applicable

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; modeled data.

Data Quality Remarks: not applicable

References: 1) U.S. Environmental Protection Agency, Office of Pollution Prevention

and Toxics and Syracuse Research Center. Copyright 2000. *EPI Suite*™, version 3.11, including MPBPWIN, version 1.41, released June 10, 2003. (Found at http://www.epa.gov/opptintr/exposure/docs/episuite.htm.)

2) Polity Industries June Meterial Sefety Pote Shoet for 2 Virgidians

2) Reilly Industries, Inc. Material Safety Data Sheet for 2-Vinylpyridine,

12 June 2000.

3) U.S. Environmental Protection Agency. Determining the Adequacy of

Existing Data; Appendix B. 10 February 1999 draft, available at

http://www.epa.gov/chemrtk/datadfin.htm.

Record Last Changed: 11/20/03

Order Number for Sorting: M1

Boiling Point

ID: 2

2-Vinylpyridine **Test Substance Identity:**

Test Substance Purity: not stated

Test Substance Remarks: not applicable

Method / Guideline Followed: not stated

GLP: not stated

Year: not stated

Test Conditions Remarks: not applicable

Boiling Point in °C: 159.5°C

Pressure: not stated

Pressure Unit: not stated **Decomposition:** not stated

Results Remarks: not applicable

Conclusions: Boiling point is adequately characterized in a reliable reference book.

Conclusions Remarks: Conclusions of the data submitter.

Klimisch Code = 2 **Data Quality Reliability:**

Reliable with restrictions; data reported in a reliable reference book.

Data Quality Remarks: not applicable

References: CRC Handbook of Chemistry and Physics; Lide, David. R., ed.

80th edition; CRC Press, Boca Raton, FL, 1999.

Supporting References:

Hawley's Condensed Chemical Dictionary; Lewis, Richard J., Sr., ed. 13th edition; John Wiley & Sons: New York, NY; 1998. (reports b.p. = 159°C)

Lange's Handbook of Chemistry; Dean, John A., ed. 14th edition; McGraw-Hill: New York, NY, 1992. (reports b.p. =

158-159°C)

11/25/03 Record Last Changed:

Order Number for Sorting:

Vapor Pressure

ID: 3

Test Substance Identity: 2-Vinylpyridine
Test Substance Purity: not applicable
Test Substance Remarks: not applicable

Method / Guideline Followed: Modeled using EPI Suite™ v3.11, MPBPWIN v1.41.

GLP: no

Year: 2003

Test Conditions Remarks: not applicable

Vapor Pressure Value: 2.57 mm Hg

Temperature (°C): 25°C

Decomposition: not applicable

Results Remarks: Modeled melting point using MPBPWIN v1.41 resulted in a mean vapor

pressure of 2.57 mm Hg (mean of Antoine & Grain Methods, which modeled 2.8 and 2.3 mm Hg, respectively). An additional method (Mackay

Method) resulted in a modeled vapor pressure of 3.48 mm Hg.

Conclusions: The modeled vapor pressure of 2.57 mm Hg fits extremely well with vapor

pressure estimations calculated from the available boiling points of 2-Vinylpyridine at reduced pressures, using the estimation method found in Lyman 1982 (see Reference #2). Using data from boiling point data from the Beilstein database (see Reference #3), the following extrapolation

calculations were performed:

BP(°C)	P (mm Hg)	702 @ 25°C (mm kg)
158.5	760	2.93
71	32	3.15
71.5	32	3.07
67.5	29	3.37
71	30	2.92
80	30	1.81
80.5	29	1.69
68.5	23	2.44
71	25	2.36
64.5	20	2.57
60	17	2.72
62	15	2.12
54	14.5	3.15
55	11	2.19
43	10	3.81
49	10	2.74
48.5	10	2.82
36	3	1.59
42	4	1.52
83	3	0.09
	Average VP (0.9396)	2.45 mm Hg

Conclusions Remarks: Conclusions of the data submitter.

Data Quality Reliability:

Klimisch Code = 2

Reliable with restrictions; modeled data.

Data Quality Remarks:

not applicable

References:

1) U.S. Environmental Protection Agency, Office of Pollution Prevention and Toxics and Syracuse Research Center. Copyright 2000. EPI Suite™, version 3.11, including MPBPWIN, version 1.41, released June 10, 2003. (Found at http://www.epa.gov/opptintr/exposure/docs/episuite.htm.) 2) Lyman, WJ; Reehl, WF; Rosenblatt, DH. 1982. Handbook of Chemical Property Estimation Methods; Environmental Behavior of Organic

Compounds; "Chapter 14: Vapor Pressure". ISBN 0-07-039175-0,

McGraw-Hill, New York, New York, USA.

3) Beilstein Database, 2003, produced by BEILSTEIN Chemiedaten GmbH, Frankfurt, Germany. Access provided by STN International, Chemical Abstracts Service, Columbus, Ohio. (Found at

http://www.cas.org.)

Record Last Changed:

11/25/03

Order Number for Sorting:

M-2

General Remarks:

Partition Coefficient

ID:

2-Vinylpyridine **Test Substance Identity:**

Test Substance Purity: not stated

Test Substance Remarks: not applicable

Method / Guideline Followed: not stated

GLP: not stated Year: not stated

not applicable **Test Conditions Remarks:**

Log Pow: log Kow = 1.54

not stated Temperature (°C):

not applicable **Results Remarks:**

Partition coefficient is adequately characterized in a reliable reference **Conclusions:**

book.

Conclusions of the data submitter. **Conclusions Remarks:**

Data Quality Reliability: Klimisch Code = 2 Reliable with restrictions; data reported in a reliable reference book.

Data Quality Remarks: not applicable

References: Chemicals Inspections and Testing Institute. 1992. Biodegradation and

Bioaccumulation Data of Existing Chemicals Based on the CSCL Japan. Japan Chemical Industry Ecology-Toxicology and Information Center. ISBN 4-89074-101-1, Japan. As referenced in Hazardous Substances Data Bank®, National Library of Medicine, Bethesda, Maryland, USA.

Found at http://toxnet.nlm.nih.gov/.

11/21/03 **Record Last Changed:**

Order Number for Sorting: 3

Water Solubility

ID: 5

Test Substance Identity: 2-Vinylpyridine

Test Substance Purity: not stated

Test Substance Remarks: not applicable

Method / Guideline Followed:

owed: not stated

Year: not stated

Test Conditions Remarks: not applicable

Value (mg/L @ °C): 2.75 x 10⁴ mg/L @ 20°C

Description of Solubility: very soluble

pH value and concentration at

temperature °C

GLP:

not stated

not stated

pKa value at 25°C: 4.98

Results Remarks: Reference reports value of 2.75 g/100 mL @ 20°C.

Conclusions: Water solubility is adequately characterized in a reliable reference source.

Conclusions Remarks: Conclusions of the data submitter.

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; data reported in a reliable reference book.

Data Quality Remarks: not applicable

References: Scriven, EFV; Toomey, JE; Murugan, R. 1996. "Pyridine and Pyridine

Derivatives" in Kirk-Othmer Encyclopedia of Chemical Technology, 4th Edition, "Volume 20, Power Generation to Recycling, Glass", John Wiley &

Sons, New York, New York, USA. p. 644.

Record Last Changed: 11/21/03

Order Number for Sorting: 4

Photodegradation

ID:

6

Test Substance Identity:

2-Vinylpyridine

Test Substance Purity:

97%

Test Substance Remarks:

Source: Aldrich Chemical

Method / Guideline Followed:

not stated

Test Type:

photodegradation study

GLP:

not stated

Year:

1993

Light Source:

"cool white" fluorescent; black lamp irradiation

Light Spectrum (nm):

not stated

Spectrum of Substance:

not stated

Test Conditions Remarks:

"Initial experiments were carried out to check that 2-vinylpyridine did not undergo photolysis under normal room fluorescent lights, black lamp irradiation, or react with NO2. Under "cool-white" fluorescent lighting at an intensity a factor of ~ 11 higher than that in an office room, < 5% decay of 2-vinylpyridine occurred over a 5.2-h period (equivalent to ~ 7 days of normal office lighting). Similarly, irradiation with black lamps at the maximum light intensity (which corresponds to an NO₂ photolysis rate of ~ 0.45 min⁻¹) for 15 min led to no observable decay of 2-vinylpyridine within

the measurement uncertainties."

Concentration of Substance:

neat

Temperature (°C):

25°C

Direct Photolysis

not stated

- Half-Life (t 1/2):

Direct Photolysis - Degradation % after: 0 after 7 day-equivalents

Direct Photolysis

- Quantum Yield:

not stated

Indirect Photolysis

ozone; OH radical; NO3 radicals

- Sensitizer (type):

Indirect Photolysis - Sensitizer Concentration: see Results Remarks

Indirect Photolysis

see Results Remarks

Indirect Photolysis

- Rate Constant:

see Results Remarks

– Degradation % after: **Breakdown Products:**

ves (see Results Remarks)

Results Remarks:

Ozone Reaction: 1.94×10^{14} molecule·cm⁻³ of 2-vinylpyridine was reacted with 1.26×10^{14} molecule·cm⁻³ of O₃. A least-squares analysis of ozone decay rates yielded a rate constant of 10⁻¹⁷ cm³ molecule⁻¹ s⁻¹. Major products observed were 2-pyridinecarboxaldehyde and formaldehyde of vields of 80 \pm 9% and 34 \pm 5%, respectively. Minor products were pyridine,

carbon dioxide, carbon monoxide and formic acid.

OH Radical Reaction: Initial experiments showed that 2-vinvlovridine reacts with gaseous nitric acid that is present in the NO2; subsequent experiments were designed using pyridine as an acid scavenger. 24 x 10¹³ molecule cm⁻³ of CH₃ONO and NO were reacted with 2.4 x 10¹ molecule·cm⁻³ of 2-vinylpyridine (and an equivalent amount of isoprene, for determining relative reaction rate) in the presence of pyridine. A leastsquares analysis of the data yielded a rate constant of 10⁻¹¹ cm³ molecule⁻¹ s¹. 2-Pyridinecarboxaldehyde was identified as the major product of the reaction, with a yield of 78 ± 14%.

Conclusions:

For ambient atmospheric conditions, the calculated lifetime of 2vinylpyridine due to the reaction with OH radicals, NO₃ radicals and O₃ is ~ 3 hours, ~ 4 hours and ~ 1 day, respectively. This assumes that 2vinylpyridine reacts with NO3 radicals with a rate constant similar to that of styrene. (Results assume ambient concentrations of the following: OH radicals, a 12-hour average of 1.6 x 10⁶ molecule cm⁻³; NO₃ radicals, a 12hour average of 5 x10⁸ molecule·cm⁻³; and O₃, a 24-hour average of 7 x 10¹¹ molecule·cm⁻³.)

It is possible that the removal of 2-vinvlovridine with gaseous nitric acid is competitive with these reactions as an atmospheric loss process.

In each case, 2-vinylpyridine was found to react to form 2-pyridinecarboxaldehyde in an analogous fashion to the breakdown of styrene to benzaldehyde, in high yield.

Conclusions Remarks:

Conclusions of the authors (see References section).

Data Quality Reliability:

Klimisch Code = 2

Reliable with restrictions; acceptable, well-documented publication/study

report which meets basic scientific principles.

Data Quality Remarks:

not applicable

References:

Tuazon, EC; Arey, J; Atkinson, R; Aschmann, SM. Gas-phase reactions of 2-vinylpyridine and styrene with OH and NO₃ radicals and O₃. Environ Sci

Technol, 27, 1832-1841 (1993).

Record Last Changed:

24 October 2003

Order Number for Sorting:

26

General Remarks:

Stability in Water

7 ID:

Test Substance Identity: 2-Vinylpyridine

Test Substance Purity: not applicable not applicable **Test Substance Remarks**

Method / Guideline Followed: estimation based on chemical principles

not applicable Test Type:

2003

GLP: not applicable

Year: **Test Conditions Remarks:** not applicable

not expected to hydrolyze at environmental pH Nominal:

Measured Value (mg/L): not applicable Degradation % at pH and °C: not applicable

Half-life (t1/2) at pH and °C: not applicable

Breakdown Products: not applicable not applicable **Results Remarks:**

Conclusions: Hydrolysis is a potentially important environmental fate pathway for a range of organic chemicals, including alkyl halides, amides, amines, carbamates,

epoxides, nitriles and esters. 2-Vinylpyridine does not contain a functional group that is susceptible to hydrolysis; in fact, alkenes are known to be generally resistant to hydrolysis. As expected, attempts to model a hydrolysis rate using the HYDROWIN™ modeling program were

unsuccessful.

Conclusions of the data submitter. **Conclusions Remarks:**

Data Quality Reliability: Klimisch Code = 2 Estimation based on generally accepted chemistry principles.

Data Quality Remarks: not applicable

References: 1) Lyman, WJ; Reehl, WF; Rosenblatt, DH. 1982. Handbook of Chemical

> Property Estimation Methods; Environmental Behavior of Organic Compounds; "Chapter 17: Rate of Hydrolysis". ISBN 0-07-039175-0,

McGraw-Hill, New York, New York, USA.

2) U.S. Environmental Protection Agency, Office of Pollution Prevention and Toxics and Syracuse Research Center. Copyright 2000. EPI Suite™. version 3.11, including HYDROWIN™, version 1.67, released June 10, 2003. (Found at http://www.epa.gov/opptintr/exposure/docs/episuite.htm.)

Record Last Changed: 11/25/03

Order Number for Sorting: M-3

Transport and Distribution (Fugacity)

ID: 8

Test Substance Identity: 2-Vinylpyridine
Test Substance Purity: not applicable
Test Substance Remarks: not applicable

Method / Guideline Followed: EPI Suite™, version 3.11, including an adapted Mackay's EQC Level III

Fugacity Model

Test Type: fugacity modeling

Year: 2003

Test Conditions Remarks: not applicable

Media: air, soil, sediment and water

Estimated Distribution and

Media Concentration:

Results from EPIWIN, v. 3.11:

Mass Amount Half-Life Emissions (hr) (percent) (kg/hr) Air 79.6 3.65 1000 Water 360 13.7 Soil 360 6.59 0 Sediment 0.0296 1.44e+003 0

```
Reaction
        Fugacity
                            Advection Reaction
                                                  Advection
         (atm)
                   (kg/hr)
                           (kg/hr)
                                       (percent)
                                                  (percent)
Air
        1.16e-011
                   947
                             49.9
                                       94.7
Water
        5.28e-013
                   1.65
                             0.86
                                        0.165
                                                   0.086
Soil
        4.4e-012
                   0.795
                             0
                                        0.0795
Sediment 4.24e-013
                   0.000891
                            3.7e-005 8.91e-005
                                                 3.7e-006
```

Persistence Time: 6.26 hr Reaction Time: 6.59 hr Advection Time: 123 hr Percent Reacted: 94.9 Percent Advected: 5.07

Half-Lives (hr), (based upon Biowin (Ultimate) and Aopwin):
 Air: 3.649

Water: 360 Soil: 360 Sediment: 1440

Biowin estimate: 2.753 (weeks

Advection Times (hr):
Air: 100
Water: 1000
Sediment: 5e+004

Results Remarks: User inputs:

Water Solubility = 27,500 mg/LVapor Pressure = 2.57 mm Hg

log Kow = 1.54

Boiling Point = 159.5°C

Melting Point = -15°C

 Emission rates to air was left as model default (1000 kg/hr); emission rates to soil and water were adjusted to 0 kg/hr, due to stringent regulation on emission of 2-Vinylpyridine via these routes.

Conclusions: Environmental transport and distribution have been adequately estimated

using accepted models.

Conclusions Remarks: Conclusions of the data submitter.

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; modeled data.

Data Quality Remarks: not applicable

References: U.S. Environmental Protection Agency, Office of Pollution Prevention and

Toxics and Syracuse Research Center. Copyright 2000. *EPI Suite™*, version 3.11, including adapted Mackay's EQC Level III Fugacity Model,

released June 10, 2003. (Found at

http://www.epa.gov/opptintr/exposure/docs/episuite.htm.)

Record Last Changed: 11/25/03

Order Number for Sorting: M-4

Biodegradation

ID: 9

Test Substance Identity: 2-Vinylpyridine

Test Substance Purity: not stated

Test Substance Remarks: not applicable

Method / Guideline Followed: aerobic biodegradation screening test

Test Type: not applicable

GLP: not stated

Year: not stated

Contact time (units): 4 weeks
Inoculum: not stated

Test Conditions Remarks: not applicable

Degradation % after time: 0% over 4 weeks

Results: not readily biodegradable

Kinetic: not stated

Breakdown Products: not stated

Results Remarks: not applicable

Conclusions: Biodegradation of 2-vinylpyridine in soil is not expected to be a major fate

process based on a single aerobic screening test showing that 2-

vinylpyridine was not biodegraded over a 4-week period.

Conclusions Remarks: Conclusions reported in reference (see below).

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; data reported in a reliable reference book.

Data Quality Remarks: not applicable

References: Chemicals Inspections and Testing Institute. 1992. Biodegradation and

Bioaccumulation Data of Existing Chemicals Based on the CSCL Japan. Japan Chemical Industry Ecology-Toxicology and Information Center. ISBN 4-89074-101-1, Japan. As referenced in *Hazardous Substances Data Bank*®, National Library of Medicine, Bethesda, Maryland, USA.

Found at http://toxnet.nlm.nih.gov/.

Record Last Changed: 11/25/03

Order Number for Sorting: 59

Biodegradation

ID: 10

2-Vinylpyridine **Test Substance Identity: Test Substance Purity:** not applicable **Test Substance Remarks:** not applicable

Modeled using EPI Suite™ v3.11, BIOWIN v4.01. Method / Guideline Followed:

Test Type:

biodegradation modeling

GLP: no 2003 Year:

Contact time (units): not applicable not applicable Inoculum: **Test Conditions Remarks:** User inputs:

> Water Solubility = 27,500 mg/L Vapor Pressure = 2.57 mm Hg

log Kow = 1.54Boiling Point = 159.5°C

Melting Point = -15°C

see Remarks Degradation % after time:

Results: see Remarks Kinetic: see Remarks **Breakdown Products:** see Remarks

Results Remarks: Linear Biodegradation Probability = 0.5429

Non-Linear Biodegradation Probability = 0.4694 MITI Linear Biodegradation Probability = 0.4173 MITI Non-Linear Biodegradation Probability = 0.4005

(NOTE: Values ≥ 0.5 indicate rapid biodegradation; values < 0.5 indicate slow biodegradation.)

Survey Model - Ultimate Biodegradation = 2.7527 (weeks to months)

Survey Model - Primary Biodegradation = 3.6773 (days to weeks)

Conclusions: Modeling data suggests that 2-vinylpyridine may not biodegrade rapidly.

Conclusions Remarks: Conclusions of the data submitter.

Data Quality Reliability: Reliable with restrictions; modeled data.

Klimisch Code = 2

Data Quality Remarks: not applicable

References: U.S. Environmental Protection Agency, Office of Pollution Prevention and Toxics and Syracuse Research Center. Copyright 2000. EPI Suite™.

version 3.11, including BIOWIN™, version 4.01, released June 10, 2003, (Found at http://www.epa.gov/opptintr/exposure/docs/episuite.htm.)

Record Last Changed: 11/26/03

Order Number for Sorting: M-5

Aquatic Toxicity - Fish

ID: 11

2-Vinylpyridine **Test Substance Identity:**

Test Substance Purity: not applicable

Test Substance Remarks: not applicable

Method / Guideline Followed:: Modeled using ECOSAR Classes for Microsoft Windows, v0.99g, April

aquatic toxicity modeling **Test Type:**

GLP: no Year: 2003

Species / Strain / Supplier: fish

Analytical Monitoring: not applicable

14 days; 96-hr; 30 days; 96-hr (salt water) **Exposure Duration:**

Statistical Methods: not applicable **Test Conditions Remarks:** User inputs:

Measured Water Solubility = 27,500 mg/L

Melting Point = -15°C Measured log Kow = 1.54

Nominal Concentrations

(mg/L):

not applicable

Measured Concentrations

(mg/L):

not applicable

Unit:

not applicable

Element Value (EC50, etc):

Predicted LC₅₀ (fish, 14-day) = 355.165 mg/L Predicted LC₅₀ (fish, 96-hr) = 210.945 mg/LPredicted LC₅₀ (fish, 96-hr, SW) = 38.687 mg/L Predicted ChV (fish, 30-day) = 25.233 mg/L

Statistical Results: not applicable

Results: see "Element Value"

Conclusion: 2-Vinylpyridine is not expected to exhibit severe toxicity to fish, as

predicted by LC₅₀ modeling data.

Conclusions Remarks: Conclusions of the data submitter.

Klimisch Code = 2 **Data Quality Reliability:**

Reliable with restrictions; modeled data.

Data Quality Remarks: not applicable

References: Cash, G; Nabholz, V. U.S. Environmental Protection Agency, Office of

Pollution Prevention and Toxics, Risk Assessment Division. Copyright 2001. ECOSAR™ Classes for Microsoft Windows, version 0.99g, released April 2001. (Found at http://www.epa.gov/oppt/newchems/21ecosar.htm.)

11/26/03 Record Last Changed:

Order Number for Sorting:

M-6

General Remarks:

Aquatic Toxicity – Fish

ID: 12

Test Substance Identity: 2-Vinylpyridine
Test Substance Purity: 95% or better

Test Substance Remarks: Test substance purchased from Aldrich Chemical or MRM Research

Chemicals Lancaster Synthesis.

Method / Guideline Followed: TETRATOX protocol (see Test Conditions Remarks)

Test Type: Tetrahymena pyriformis toxicity

GLP: not stated
Year: 2001

Analytical Monitoring: not stated

Species / Strain / Supplier: Tetrahymena pyriformis, strain GL-C

Test Details: Endpoint (population density) of static 40-hour assay was measured

spectrophotometrically at 540 nm. Stock solutions were prepared in dimethyl sulfoxide, which has NOEC of 7500 mg/L; care was taken to

ensure that this level was not exceeded.

Two controls were used -- one with *T. pyriformis* without test chemicals or solvents, and the other control as a blank, containing neither test

chemical, solvent, nor ciliates.

Test conditions allowed for 8-9 cell cycles in control cultures. Each definitive test replicate consisted of 6 - 8 different concentrations with

duplicate flasks of each concentration.

Exposure Duration: 40 hours

Statistical Methods: Probit Analysis procedure of Statistical Analysis System (SAS) software

Test Conditions Remarks: TETRATOX protocol is defined in the following citation: Schultz, T.W.

1997. TETRATOX: Tetrahymena pyriformis population growth impairment endpoint-A surrogate for fish lethality. Toxicol. Methods 7:

289-309.

Nominal Concentrations (mg/L): not stated

Measured Concentrations

(mg/L):

not performed

Unit: not stated

Element Value (EC50, etc.) 50% growth inhibition concentration (IGC₅₀) = 0.57 mg/L

Statistical Results: $\log(IGC_{50}^{-1}) = 0.24$

Results Remark: While the study report focuses on establishing a QSAR model, it is

important to note that 2-vinylpyridine data were experimentally derived in

this study to compare to predicted values.

Conclusions: Authors conclude that QSAR's previously established for the estimation

of toxicity of benzene derivatives can be extended to include pyridines, with the understanding that pyridines substituted with electron-releasing groups in the ortho position may not fit the model as well. The pyridine response-surface has more that 5% more unexplained error than the benzene surface, but the response-surfaces derived for the two chemical

groups are virtually identical.

Conclusions Remarks: Conclusions of the authors (see References section).

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; acceptable, well-documented publication /

study report which meets basic scientific principles.

Data Quality Remarks: not applicable

References: Seward, JR; Cronin, MTD; Schultz, TW. Structure-toxicity analyses of

Tetrahymena pyriformis exposed to pyridines - an examination into extension of surface-response domains. 2001. SAR and QSAR in

Environmental Research, 11, 489-512.

Record Last Changed: 10/27/03

Order Number for Sorting: 48

Aquatic Toxicity - Invertebrate

ID: 13

Test Substance Identity: 2-Vinylpyridine
Test Substance Purity: not applicable
Test Substance Remarks: not applicable

Method / Guideline Followed: Modeled using ECOSAR Classes for Microsoft Windows, v0.99g, April

2001.

Test Type: aquatic toxicity modeling

GLP: no Year: 2003

Analytical Monitoring: not applicable

Species / Strain / Supplier: Daphnia magna

Test Details: not applicable

Exposure Duration: 48-hr; 16-day

Statistical Methods: not applicable

Test Conditions Remarks: User inputs:

Measured Water Solubility = 27,500 mg/L

Melting Point = -15°C
 Measured log Kow = 1.54

Nominal Concentration (mg/L): not applicable

Measured Concentration (mg/L): not applicable

Unit: not applicable

Element Value (EC50, etc.)

Jilli. Hot applicable

Predicted EC₅₀ (16-day, Daphnia) = 9.183 mg/L

Predicted LC₅₀ (48-hr, *Daphnia*) = 218.962 mg/L

Statistical Results: not applicable

Results Remarks: not applicable

Conclusions: 2-Vinylpyridine is not expected to exhibit severe toxicity to aquatic

invertebrates, as predicted by LC50 modeling data.

Conclusions Remarks: Conclusions of the data submitter.

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; modeled data.

Data Quality Remarks: not applicable

References: Cash, G; Nabholz, V. U.S. Environmental Protection Agency, Office of

Pollution Prevention and Toxics, Risk Assessment Division. Copyright 2001. *ECOSAR™ Classes for Microsoft Windows*, version 0.99g,

released April 2001. (Found at

http://www.epa.gov/oppt/newchems/21ecosar.htm.)

Record Last Changed: 11/26/03

Order Number for Sorting: M-7

Aquatic Toxicity - Aquatic Plant

ID: 14

Test Substance Identity: 2-Vinylpyridine
Test Substance Purity: not applicable
Test Substance Remarks: not applicable

Method / Guideline Followed: Modeled using ECOSAR™ Classes for Microsoft Windows, v0.99g, April

2001.

Test Type: modeled aquatic toxicity

GLP: no Year: 2003

Species / Strain / Supplier: green algae

Element Basis: not applicable

Exposure Duration: 96-hr

Analytical Monitoring: not applicable

Statistical Methods: not applicable

Test Conditions Remarks: User inputs:

Measured Water Solubility = 27,500 mg/L

Melting Point = -15°C
Measured log Kow = 1.54

Nominal Concentrations

(mg/L):

not applicable

Measured Concentrations

(mg/L):

not applicable

Unit: not applicable

Element Value (EC50, etc.): Predicted EC₅₀ (96-hr, green algae) = 133.312 mg/L

Predicted ChV (96-hr, green algae) = 10.219 mg/L

NOEC, LOEC or NOEL, LOEL:

EL: not applicable

Satisfactory Control

Response:

not applicable

Statistical Results: not applicable
Results Remarks: not applicable

Conclusions: 2-Vinylpyridine is not expected to exhibit severe toxicity to aquatic plants,

as predicted by EC₅₀ modeling data.

Conclusions Remarks: Conclusions of the data submitter.

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; modeled data.

Data Quality Remarks: not applicable

References: Cash, G; Nabholz, V. U.S. Environmental Protection Agency, Office of

Pollution Prevention and Toxics, Risk Assessment Division. Copyright 2001. *ECOSAR™ Classes for Microsoft Windows*, version 0.99g, released April 2001. (Found at http://www.epa.gov/oppt/newchems/21ecosar.htm.)

Record Last Changed: 11/26/03
Order Number for Sorting: M-8

ID:

15

Test Substance Identity:

2-Vinylpyridine

Test Substance Purity:

98.7%

Test Substance Remarks:

Lot number 20116AC -- characterization data on file with test sponsor

Method / Guideline Followed:

TSCA method: 40 CFR 789.1100, July 1989

Test Type:

acute dermal toxicity

GLP:

yes

Year:

1992

Species / Strain / Supplier:

New Zealand White rabbits (Oryctolagus cuniculus), 10-12 weeks old,

Eastern Rabbit Breeding Laboratory, Taunton, MA

Sex:

male and female

No of Animals/Sex/Dose:

5

Vehicle:

none (undiluted test substance)

Route of Administration:

dermal application

Test Conditions Remarks:

Site of application was not abraded intentionally or accidentally during preparation. Exposure period = 24 hours; observation period = 14 days. Test substance was introduced under gauze patches two single layers thick and applied directly to the skin (approximately 10% of the body surface). Gauze was moistened with USP Water for Injection and patches were secured in place by wrapping the entire trunk of the animal with

impervious bandaging.

Value (LD₅₀, etc) with Confidence Limits: $LD_{50} = 0.64 \text{ g/kg}$

Number of Deaths at Each

Dose Level:

0.90 g/kg = 10/10 died (within first hour following dosing)
0.65 g/kg = 8/10 died (within 3 hours following dosing)
0.40 g/kg = 0/10 died over 14 day observation period

Results Remarks:

Necrosis of the skin was observed at all dosing sites.

Estimated LD₅₀ was determined utilizing the "Litchfield and Wilcoxon II" program (Tallarida, RJ; Murray, RB. 1986. *Manual of Pharmacologic Calculations with Computer Programs*. New York: Springer-Verlag, pp.

159-164).

Conclusions:

Based upon the mortality and the criteria of the study protocol, the estimated LD₅₀ of the test substance has been determined to be 0.64 g/kg.

Conclusions Remarks:

Conclusions of the authors (see References section).

Data Quality Reliability:

Klimisch Code = 1

Reliable without restriction; guideline study.

Data Quality Remarks:

not applicable

References:

Fitzgerald, G. B. 1994. Acute dermal toxicity study (single exposure), amended report. Report number 92G-0361. Toxikon Corp., Woburn, MA.

Record Last Changed:

10/27/03

Order Number for Sorting:

29

General Remarks:

ID: 16

Test Substance Identity: 2-Vinylpyridine

Test Substance Purity:

97.689%

Test Substance Remarks:

Test substance obtained from W.P. Hamilton, Camden, South Carolina.

Method / Guideline Followed:

not stated (see Remarks)

Test Type:

acute dermal toxicity

GLP:

not stated

Year:

1981

Species / Strain / Supplier:

New Zealand White rabbits

Sex:

male

No of Animals/Sex/Dose:

1 animal/dose; 6 doses

Vehicle:

none (undiluted test substance)

Route of Administration:

dermal application

Test Conditions Remarks:

While no method is stated, the procedure listed is significantly similar to US EPA's harmonized OPPTS guidance for Acute Dermal Toxicity, but with fewer animals. Exposure period = 24 hours; observation period = 14 days. Volumes of test material applied ranged from 1.66 to 0.17 mL. Test substance was applied to the trunk of each rabbit under two 12 ply gauze pads, and the trunk of each animal was wrapped with a layer of Saran® Wrap, Kling® gauze bandage and Elastoplast® adhesive bandage.

Value (LD₅₀, etc) with Confidence Limits:

approximate lethal dose = 300 mg/kg

Number of Deaths at Each

Dose Level:

670 mg/kg: 1/1 died within 1 day of dosing; 450 mg/kg: 1/1 died within 2.5 hrs. of dosing; 300 mg/kg: 1/1 died within 2 hrs of dosing;

200, 90 & 60 mg/kg: 0/1 died over 14-day observation period after dosing.

Results Remarks:

not applicable

Conclusions:

2-Vinylpyridine is considered to be moderately toxic by skin absorption in male rabbits. Approximate lethal dose (ALD) is 300 mg/kg of body weight. Clinical signs included lethargy, prostration, labored breathing, aggressive behavior, convulsion, severe skin irritation and weight loss. All deaths

occurred within 1 day after dosing.

Conclusions Remarks:

Conclusions of the authors (see References section).

Data Quality Reliability:

Klimisch Code = 2

Reliable with restrictions; basic data given: comparable to

guidelines/standards with acceptable restrictions.

Data Quality Remarks:

not applicable

References:

Henry, JE. 1981. Rabbit skin absorption (ALD) with pyridine, 2-ethenyl-, with cover letter. Haskell Laboratories. As submitted to USEPA under TSCA §8(e) Compliance Audit Program agreement by DuPont Chemical,

1992, OTS#0571402.

Record Last Changed: 11/3/03
Order Number for Sorting: 36

ID: 17

Test Substance Identity: 2-Vinylpyridine

Test Substance Purity: 98.95%

Test Substance Remarks: not applicable

Method / Guideline Followed: not stated

Test Type: acute oral toxicity

GLP: yes
Year: 1983
Species / Strain / Supplier: rat

Sex: male and female

No of Animals/Sex/Dose: not stated

Vehicle: none (undiluted test substance)

Route of Administration: oral

Test Conditions Remarks: No data available on dose levels, fasting or length of post observation

period.

Value (LD₅₀, etc) with Confidence Limits:

Oral LD₅₀ (rat) = 336 mg/kg (confidence limits = 240 - 472 mg/kg)

Number of Deaths at Each

Dose Level:

not stated

Results Remarks: not applicable

Conclusions: Oral LD₅o in male and female rats was 336 mg/kg. Clinical signs of toxicity

included prostration, weakness, tremors, vasodilatation, excessive

salivation and anorexia.

Conclusions Remarks: Conclusions of the authors (see References section).

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; basic data given: comparable to guidelines /

standards.

Data Quality Remarks: not applicable

References: Anonymous. 1983. Basic Toxicity of 2-Vinylpyridine. Eastman Kodak

Company, Corporate Health and Environment Laboratories. As submitted to USEPA under TSCA §8(e) Compliance Audit Program agreement by

Eastman Kodak Company, 1992, OTS#0546362.

Record Last Changed: 11/3/03

Order Number for Sorting: 17

ID: 18

Test Substance Identity: 2-Vinylpyridine

Test Substance Purity: 98.95%

Test Substance Remarks: not applicable

Method / Guideline Followed: not stated

Test Type: acute oral toxicity

GLP: yes
Year: 1983
Species / Strain / Supplier: rat

Sex: male and female

No of Animals/Sex/Dose: not stated

Vehicle: 20% compound in corn oil

Route of Administration: oral

Test Conditions Remarks: No data available on dose levels, length of fasting period, or length of post-

dosing observation period. No data available on dosing formulation

preparation or dose volumes administered.

Value (LD₅₀, etc) with Confidence Limits:

Fasted rats: $LD_{50} = 951$ mg/kg for males (C.I. = 677 - 1336 mg/kg);

 $LD_{50} = 673 \text{ mg/kg}$ for females (C.I. = 479 - 945 mg/kg).

Fed rats: $LD_{50} = 951 \text{ mg/kg}$ for both males and females (C.I. = 677 -

1336 mg/kg).

Number of Deaths at Each

Dose Level:

not stated

Results Remarks: not applicable

Conclusions: Oral LD₅₀ of 20% solution in corn oil was 951 mg/kg in fasted male rats and

673 mg/kg in fasted female rats. In fed rats, LD $_{50}$ of the 20% solution was 951 mg/kg for both males and females. Clinical signs of toxicity included

prostration, weakness, tremors, diarrhea and vasodilatation.

Conclusions Remarks: Conclusions of the authors (see References section).

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; basic data given: comparable to guidelines /

standards.

Data Quality Remarks: not applicable

References: Anonymous. 1983. Basic Toxicity of 2-Vinylpyridine. Eastman Kodak

Company, Corporate Health and Environment Laboratories. As submitted to USEPA under TSCA §8(e) Compliance Audit Program agreement by

Eastman Kodak Company, 1992, OTS#0546362.

Record Last Changed: 11/3/03

Order Number for Sorting: 17

ID:

19

Test Substance Identity:

2-Vinylpyridine

Test Substance Purity:

98.95%

Test Substance Remarks:

not applicable

Method / Guideline Followed:

not stated

Test Type:

24-hr occluded skin irritation

GLP:

yes

Year:

1983

Species / Strain / Supplier:

guinea pig

Sex:

not stated

No of Animals/Sex/Dose:

10 animals / dose

Vehicle:

none (undiluted test substance)

Route of Administration:

dermal

Test Conditions Remarks:

Dose levels = 5, 2, 1, 0.5, 0.35, 0.2, 0.1, 0.05 mL/kg. No data on observation period or application site preparation. Scoring based on qualitative parameters of "slight, moderate, strong or severe".

Value (LD₅₀, etc) with Confidence Limits:

strong dermal irritant

Number of Deaths at Each

Dose Level:

All guinea pigs given 5, 2, 1, 0.5 or 0.35 mL/kg died, six of ten guinea pigs given 0.2 mL/kg died, single guinea pig given 0.1 or 0.05 mL/kg died.

Results Remarks:

"questionable" for estimated corrosivity

Conclusions:

Skin irritation was strong as determined by a standardized 24-hour

occluded skin irritation test in guinea pigs.

Conclusions Remarks:

Conclusions of the authors (see References section).

Data Quality Reliability:

Klimisch Code = 2

Reliable with restrictions; basic data given: comparable to guidelines /

standards.

Data Quality Remarks:

not applicable

References:

Anonymous. 1983. Basic Toxicity of 2-Vinylpyridine. Eastman Kodak Company, Corporate Health and Environment Laboratories. As submitted to USEPA under TSCA §8(e) Compliance Audit Program agreement by

Eastman Kodak Company, 1992, OTS#0546362.

Record Last Changed:

11/3/03

Order Number for Sorting:

17

General Remarks:

ID: 20

Test Substance Identity:

2-Vinylpyridine

Test Substance Purity:

98.95%

Test Substance Remarks:

not applicable

Method / Guideline Followed:

not stated

Test Type:

skin irritation (open, repeated dose)

GLP:

yes

Year:

1983

Species / Strain / Supplier:

guinea pig

Sex:

not stated

No of Animals/Sex/Dose:

5 animals/dose

Vehicle:

none (undiluted test substance)

Route of Administration:

dermal

Test Conditions Remarks:

Test involved 4 to 7 doses of 0.1 mL/kg each. No data on length of observation period or site preparation. Qualitative scoring only.

Value (LD₅₀, etc) with Confidence Limits:

Strong exacerbation; skin absorption noted. Dermal LD₅₀ is noted as 0.16

mL/kg, but not test details are reported for the LD₅₀ procedure.

Number of Deaths at Each

Dose Level:

0.1 mL/kg killed all guinea pigs after 4-7 doses.

Results Remarks:

not applicable

Conclusions:

The dermal LD_{50} was 0.16 mL/kg. There was evidence of percutaneous

absorption.

Conclusions Remarks:

Conclusions of the authors (see References section).

Data Quality Reliability:

Klimisch Code = 2

Reliable with restrictions; basic data given: comparable to guidelines /

standards.

Data Quality Remarks:

not applicable

References:

Anonymous. 1983. Basic Toxicity of 2-Vinylpyridine. Eastman Kodak Company, Corporate Health and Environment Laboratories. As submitted to USEPA under TSCA §8(e) Compliance Audit Program agreement by

Eastman Kodak Company, 1992, OTS#0546362.

Record Last Changed:

11/3/03

Order Number for Sorting:

17

General Remarks:

ID: 21

2-Vinylpyridine **Test Substance Identity:**

Test Substance Purity:

98.95%

Test Substance Remarks:

not applicable

Method / Guideline Followed:

not stated

Test Type:

skin sensitization

GLP:

yes

Year:

1983

Species / Strain / Supplier:

guinea pig

not stated

No of Animals/Sex/Dose:

10 animals total

Vehicle:

none (undiluted test substance)

Route of Administration:

dermal

Test Conditions Remarks:

Report indicates "standardized skin sensitization" test, but no details are available on study design. No details on induction or challenge phases or on concentration selection. No details on grading system used or on

positive/negative controls.

Value (LD50, etc) with **Confidence Limits:**

Estimated to be a moderate risk for skin sensitization in humans.

Number of Deaths at Each

Dose Level:

not stated

Results Remarks:

Sensitization responses:

2/10 animals showed no response; 3/10 showed weak response; 4/10 showed moderate response; 1/10 showed potent response.

Conclusions:

In a standardized skin sensitization test in guinea pigs, this compound elicited reactions indicating an estimated moderate risk for human

sensitization.

Conclusions Remarks:

Conclusions of the authors (see References section).

Data Quality Reliability:

Klimisch Code = 2

Reliable with restrictions; basic data given: comparable to guidelines /

standards.

Data Quality Remarks:

not applicable

References:

Anonymous. 1983. Basic Toxicity of 2-Vinylpyridine. Eastman Kodak Company, Corporate Health and Environment Laboratories. As submitted to USEPA under TSCA §8(e) Compliance Audit Program agreement by

Eastman Kodak Company, 1992, OTS#0546362.

Record Last Changed:

11/3/03

Order Number for Sorting:

17

General Remarks:

ID: 22

Test Substance Identity: 2-Vinylpyridine

Test Substance Purity: 98.95%

Test Substance Remarks: not applicable

Method / Guideline Followed: not stated

Test Type: eye irritation

GLP: yes
Year: 1983
Species / Strain / Supplier: rabbit

Sex: not stated

No of Animals/Sex/Dose: 3 animals per test (washed and unwashed eyes)

Vehicle: none (undiluted test substance)

Route of Administration: ocular

Test Conditions Remarks: not applicable

Value (LD₅₀, etc) with Confidence Limits:

Strong eye irritant in unwashed eyes; moderate eye irritant in washed eyes.

Number of Deaths at Each

Dose Level:

not stated

Results Remarks: Unwashed eyes: 3/3 showed strong irritation.

Washed eyes: 3/3 showed moderate irritation.

Corneal and adnexal staining in all washed and unwashed eyes. Prompt

irrigation with distilled water was palliative.

Conclusions: Rabbit eye irritation was strong in unwashed (3/3) and moderate (3/3) in

washed eyes; there was corneal and adnexal staining in all washed and unwashed eyes. Prompt irrigation with distilled water was palliative.

Conclusions Remarks: Conclusions of the authors (see References section).

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; basic data given: comparable to guidelines /

standards.

Data Quality Remarks: not applicable

References: Anonymous. Basic Toxicity of 2-Vinylpyridine. Eastman Kodak Company,

Corporate Health and Environment Laboratories, 1983. As submitted to USEPA under TSCA §8(e) Compliance Audit Program agreement by

Eastman Kodak Company, 1992, OTS#0546362.

Record Last Changed: 11/3/03
Order Number for Sorting: 17

ID: 23

Test Substance Identity: 2-Vinylpyridine
Test Substance Purity: not stated

Test Substance Remarks: Source of material: Reilly Industries, Inc., lot #40310AA.

Method / Guideline Followed: 49 CFR 173.136-137, as of 16 May 1994

Test Type: DOT skin corrosion

GLP: no **Year**: 1994

Species / Strain / Supplier: New Zealand albino rabbit

Sex: 5 males; 1 female

No of Animals/Sex/Dose: all animals received 0.5 mL applied to intact skin free from hair

Vehicle: none (undiluted test substance)

Route of Administration: dermal

Test Conditions Remarks: Test sites were immediately covered with an adhesive-backed gauze

patch, secured with 3" non-irritating Durapore tape. After one hour of exposure, patches were removed and test sites wiped clean to prevent further exposure. Test sites were evaluated at 1 hour and 48 hours post-dosing. Tissue destruction was considered to have occurred if there was ulceration and/or necrosis. Epidermal sloughing, erythema, edema or fissuring were not considered to be tissue destruction. Results reported

simply as positive or negative for tissue destruction.

Value (LD₅₀, etc) with Confidence Limits:

Visible necrosis of the skin tissue was observed at all test sites, 48 hours

after dosing.

Number of Deaths at Each

Dose Level:

none

Results Remarks: not applicable

Conclusions: Visible necrosis of the skin tissue was observed at all test sites, 48 hours

after dosing. These results place the test material into Class I, Packing

Group II.

Conclusions Remarks: Conclusions of the authors (see References section).

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; basic data given: comparable to guidelines /

standards.

Data Quality Remarks: not applicable

References: Barr, K; Wnorowski, G. 1994. DOT Skin Corrosion, Study #T-2927,

Product Safety Labs, East Brunswick, NJ, unpublished.

Record Last Changed: 12/8/03

Order Number for Sorting: 40

Genetic Toxicity - In Vitro

ID: 24

Test Substance Identity:

2-Vinylpyridine

Test Substance Purity:

> 98%

Test Substance Remarks:

Chemicals for the study were purchased from commercial sources.

Method / Guideline Followed:

Ames test

Test Type:

reverse mutation assay

Testing System:

bacterial

GLP:

not stated

Year:

1980

Species / Strain or Cell Type:

Salmonella typhimurium; strains TA1535, TA98 and TA100

Metabolic Activation:

both with and without S9 (rat liver induced with Aroclor 1254)

Concentrations Tested:

0, 0.1 and 0.5 mL/9 L desiccator

Statistical Methods:

not stated

Test Conditions Remarks:

Triplicate plates prepared for each chemical concentration, both with and without S9 mix. Negative and positive controls were run with every experiment (no details given). Experiments were repeated if results were

equivocal.

Dosing via exposure in sealed desiccators for 7 hours, followed by incubation (37°C) for 40-50 hours. Plates with S9 mix were in different

desiccators from plates without S9 mix.

Result:

negative

Cytotoxic Concentration:

Severe toxicity observed at 0.5 mL/9 L concentration level, invalidating

results at this level.

Genotoxic Effects:

At 0.1 mL/9 L concentration, no mutagenic effects were observed, both

with and without metabolic activation.

Statistical Results:

not stated

Results Remarks:

No mutagenic effects observed in concentrations tested, both with and

without metabolic activation.

Conclusions:

2-Vinylpyridine was not observed to exhibit mutagenicity in this assay.

Mutagenic activity could not be adequately assessed at high

concentrations because of severe toxicity.

Conclusions Remarks:

Conclusions of the authors (see References section).

Data Quality Reliability:

Klimisch Code = 1

Comparable to guideline study.

Data Quality Remarks:

not applicable

References:

Simmon, VF; Baden, JM. Mutagenic activity of vinyl compounds and

derived epoxides. 1980. Mutation Research, 78, 227-231.

Record Last Changed:

10/29/03

Order Number for Sorting:

11

General Remarks:

Genetic Toxicity - In Vitro

25 ID:

2-Vinylpyridine **Test Substance Identity:**

98.3% **Test Substance Purity:**

Upon completion of the test, analysis of the residual test substance **Test Substance Remarks:**

revealed no stability problem.

Method / Guideline Followed: "Guidelines for Screening Mutagenicity Testing of Chemicals", Japan

mammalian cell chromosomal aberration assay **Test Type:**

Testing System: mammalian cells

GLP: yes

not stated (after 1991, per test program history) Year:

Chinese hamster adenofibroblast cell strain; obtained from National Species / Strain or Cell Type:

Sanitation Test Center, 15 Nov 1984

both with and without S9 (rat liver induced with phenobarbital and 5,6-**Metabolic Activation:**

benzoflavone)

term tests).

Concentrations Tested: maximum 30.0 μg/mL (24-hr continuous); 15.0 μg/mL (48-hr continuous);

120 µg/mL (short-term, no S9); 300 µg/mL (short-term, with S9)

Statistical Methods:

Continuous treatment: test substance solution administered continuously **Test Conditions Remarks:**

either for 24 hours or 48 hours.

Short-term treatment: test substance solution administered for 6 hours either with or without S9 mix, then transferred to fresh culture medium and

cultivated an additional 18 hours.

Positive controls were mitomycin C (0.05 µg/mL for 24-hr treatment: 0.025 μg/mL for 48-hr treatment) and cyclophosphamide (12.5 μg/mL for short-

2 plates per test; solvent was DMSO.

Concentrations (in µg/mL):

24-hr continuous: 30.0, 15.0, 7.5, 3.75, 0 48-hr continuous: 15.0, 7.5, 3.75, 1.88, 0 Short-term, -S9: 120, 60, 30, 15, 0

Short-term, +S9: 300, 150, 75, 37.5, 0

Result: positive

50% cell proliferation inhibition concentrations (by Probit method) = **Cytotoxic Concentration:**

> 33.4 µg/mL (24-hr continuous); 7.90 µg/mL (48-hr continuous); 109 μg/mL (short-term, without S9); 147 µg/mL (short-term, with S9)

Clear dose-dependent induction of chromosomal structural aberration was **Genotoxic Effects:**

observed in all test systems. D20 value = 0.00557 mg/mL; TR value =

2,000.

No notable changes such as precipitation were observed during the test.

Statistical Results: For test systems exhibiting positive results, the D20 value was computed

by the least squares method, and the TR value was computed by dividing

the frequency of occurrence (%) of chromatid exchange at the

corresponding dose by the test dose (mg/mL).

Results Remarks: Positive for clastogenicity, both with and without metabolic activation.

Negative for polyploidy. Lowest concentration producing cytogenetic

effects in vitro: 0.00557 mg/mL over 24-hrs (clastogenicity)

Chromosomal analysis conducted according to classification methods of the Mammal Test Section, Japan Environmental Mutagen Society.

No polyploid cell inducing effect was found in any of the treatment groups.

Conclusions: On the basis of the above test results, 2-vinylpyridine was determined to

have a positive effect on the inducement of chromosomal aberrations in

cultivated mammal cells under the conditions of this test.

Conclusions Remarks: Conclusions of the authors (see References section).

Data Quality Reliability: Klimisch Code = 1

Guideline study: "Guidelines for Screening Mutagenicity Testing of

Chemicals", Japan

Data Quality Remarks: not applicable

References: Nakajima, Madoka, et al. In vitro chromosomal aberration test of 2-

vinylpyirdine on cultured Chinese hamster cells. Biosafety Research Center, Foods, Drugs and Pesticides (An-pyo Center), Japan, 582-2 Shioshinden Arahama, Fukude-cho, Iwata-gun, Shizuoka, 437-12, Japan. (Located at Japan's Global Information Network on Chemicals, found at

http://wwwdb.mhlw.go.jp/ginc/index.html.)

Record Last Changed: 10/30/03

Order Number for Sorting: 55

Genetic Toxicity - In Vitro

ID: 26

Test Substance Identity: 2-Vinylpyridine

Test Substance Purity: 98.3%

Test Substance Remarks: Upon completion of the test, analysis of the residual test substance

revealed no stability problem.

Method / Guideline Followed: "Guidelines for Screening Mutagenicity Testing of Chemicals", Japan

Test Type: reverse mutation assay

Testing System: bacterial

Species / Strain or Cell Type:

GLP: yes

Year: not stated (after 1991, per test program history)

coli WP2 uvrA

Metabolic Activation: both with and without S9 (rat liver induced with phenobarbital and 5,6-

benzoflavone)

Concentrations Tested: without S9: 39.1 to 2500 µg/plate; with S9: 156 - 5000 µg/plate

Statistical Methods: not stated

Test Conditions Remarks: Source of Salmonella typhimurium: Ames, University of California, 9

September 1985. Source of Escherichia coli: National Sanitation Test

Salmonella typhimurium TA100, TA98, TA1535, TA1537 and Escherichia

Center, 16 March 1985.

Positive control substances included 2-(2-furyl)-3-(5-nitro-2-

furyl)acrylamide, sodium azide, 9-aminoacrylidine, 2-aminoanthracene. Negative controls (no test substance) were also run throughout the

experiment.

Three plates/test. Solvent: DMSO. Each test was independently

conducted twice.

Result: negative in *S. typhimurium*; positive in *E. coli* (+S9 only)

Cytotoxic Concentration: Both the +S9 and -S9 groups exhibited growth-inhibiting activity due to 2-

vinylpyridine treatment at high doses.

Genotoxic Effects: None observed with *S. typhimurium* strains.

E. coli WP2 uvrA strain to which S9 mix was added showed a clear increase accompanying dose-effect correlation. Reproducibility was

confirmed. Relative mutation activity of 10.7 was exhibited.

No genotoxic effects were observed in E.coli WP2 uvrA strain without

metabolic activation.

Statistical Results: not conducted

Results Remarks: Negative for genetic effects with S. typhimurium (all strains), both with and

without metabolic activation. Clear positive mutagenic response in E. coli

strain with metabolic activation only (negative without activation).

No notable changes such as precipitation were observed during the test.

Conclusions: No genetic effects were observed with any Salmonella typhimurium strain,

both with and without metabolic activation.

A clear positive mutagenic response was obtained in *E. coli* WP2 uvrA with metabolic activation only (negative response without metabolic activation).

Conclusions Remarks: Conclusions of the authors (see References section).

Data Quality Reliability:

Klimisch Code = 1

Guideline study: "Guidelines for Screening Mutagenicity Testing of

Chemicals", Japan

Data Quality Remarks:

not applicable

References:

Nakajima, Madoka, et al. Reverse mutation test of 2-vinylpyirdine on bacteria. Biosafety Research Center, Foods, Drugs and Pesticides (Anpyo Center), Japan, 582-2 Shioshinden Arahama, Fukude-cho, Iwata-gun, Shizuoka, 437-12, Japan. (Located at Japan's Global Information Network on Chemicals, found at http://wwwdb.mhlw.go.jp/ginc/index.html.)

Record Last Changed:

10/30/03

Order Number for Sorting:

56

General Remarks:

Genetic Toxicity - In Vitro

ID: 27

Test Substance Identity 2-Vinylpyridine
Test Substance Purity not stated

Test Substance Remarks

Test substance was obtained from Aldrich Chemical Company.

Method / Guideline Followed: Adapted from procedure described by Borenfreund and Puerner (1984)

Test Type: mammalian cell cytotoxicity assay (neutral red)

Testing System: mammalian cells

GLP: not stated
Year: 1995

Species / Strain or Cell Type: WB rat liver cells, JB-6 (clone 25) mouse keratinocytes, CHO (K1) cells,

A549 human lung carcinoma cells, CCD-11 Lu human lung fibroblasts

Metabolic Activation: none

Concentrations Tested: not stated

Statistical Methods: linear regression

Test Conditions Remarks: WB rat liver cells and mouse keratinocytes were obtained from Michigan

State University. All other cells obtained through American Type Culture Collection. Ammonium hydroxide and n-butanol were run as positive controls. Exposures were at 37°C incubation for 24 hours. Absorbance was measured at 540 nm, and concentration vs. absorbance was plotted. EC₅₀ was defined as the chemical concentration required to reduce the absorbance value by 50% with respect to the solvent or untreated controls.

Result: no result (toxic to test system)

Cytotoxic Concentration: $EC_{50} = 0.5 \text{ mM}$ in Chinese hamster ovary cells

Genotoxic Effects: not evaluated

Statistical Results: linear regression

Results Remarks: Cytotoxicity observed (see "Cytotoxic Concentration" below).

Conclusions: 2-Vinylpyridine caused more cytotoxicity to CHO cells than did pyridine or

4-methylpyridine, but was an order of magnitude less cytotoxic than 4-vinylpyridine. Authors suggest that the increased cytotoxicity of vinylpyridines may be due to the reactivity of the vinyl group to cellular

proteins and DNA.

Conclusions Remarks: Conclusions of the authors (see References section).

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; acceptable, well-documented publication/study

report which meets basic scientific principles.

Data Quality Remarks: not applicable

References: Bombick, DW; Doolittle, DJ. 1995. The role of chemical structure and cell

type in the cytotoxicity of low-molecular-weight aldehydes and pyridines. In

Vitro Toxicology, 8(4), 349-356.

Record Last Changed: 10/30/03

Order Number for Sorting: 32

Genetic Toxicity - In Vitro

ID:

Test Substance Identity: 2-Vinylpyridine

> 99% **Test Substance Purity:**

Test substance obtained from Aldrich Chemical and further distilled. **Test Substance Remarks:**

Method / Guideline Followed: Ames test

reverse mutation assay Test Type:

Testing System: bacterial not stated GLP: 1992 Year:

Salmonella typhimurium; strains TA1535, TA 1538, TA98 and TA100 Species / Strain or Cell Type:

Metabolic Activation: both with and without S9 (rat liver induced with Aroclor)

5, 10, 25 and 50 µmol/plate **Concentrations Tested:**

Statistical Methods: not stated **Test Conditions Remarks:** not applicable negative Result:

At 5 and 10 µmol/plate, survival rates were 80-92%. **Cytotoxic Concentration:**

At 25 and 50 µmol/plate, survival rates were 30-65%.

Not mutagenic in concentrations tested. Genotoxic Effects:

Statistical Results: not stated

No mutagenic effects observed in concentrations tested, both with and Results Remarks:

without metabolic activation.

Conclusions: 2-Vinylpyridine was not mutagenic in concentrations of 5, 10, 25 and 50

µmol/plate in Ames tester strains TA1535, TA 1538, TA98 and TA100 without or with metabolic activation by Aroclor-induced S9 fraction.

Conclusions of the authors (see References section). **Conclusions Remarks:**

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; basic data given: comparable to

guidelines/standards.

not applicable **Data Quality Remarks:**

References: Brunnemann, KD; Rivenson, A; Cheng, SC; Saa, V; Hoffmann, D. 1992. A

study of tobacco carcinogenesis XLVII. Bioassays of vinylpyridines for genotoxicity and for tumorigenicity in A/J mice. Cancer Letters, 65, 107-

113.

11/7/03 **Record Last Changed:** 34

Order Number for Sorting:

General Remarks: not applicable

Genetic Toxicity - In Vitro

ID:

29

Test Substance Identity:

2-Vinylpyridine

Test Substance Purity:

> 99%

Test Substance Remarks:

Test substance obtained from Aldrich Chemical and further distilled.

Method / Guideline Followed:

hepatocyte primary culture / DNA repair test according to G. M. Williams

(see "Test Conditions Remarks").

Test Type:

rat hepatocyte DNA repair test

Testing System:

mammalian cell

GLP:

not stated

Year:

1992

Species / Strain or Cell Type:

rat hepatocytes

Metabolic Activation:

none

Concentrations Tested:

2.5, 5.0, 7.5, 10.0 mmol

Statistical Methods:

not stated

Test Conditions Remarks:

Williams test method referenced at *Mutat. Res.*, 1989, 221, 263-286. Similar to OPPTS 870.5550. 4-(Methylnitrosamino)-1-(3-pyridyl)-1-

butanone (NNK) was used as a positive control.

Result:

negative

Cytotoxic Concentration:

2-Vinylpyridine was toxic to rat hepatocytes at 5,0, 7.5 and 10.0 mmol

doses. (Toxicity defined as < 80% viable liver cells.)

Genotoxic Effects:

2-Vinylpyridine did not exhibit genotoxicity at 2.5 mmol dose.

Statistical Results:

not stated

Results Remarks:

No genotoxicity observed at non-toxic doses.

Conclusions:

2-Vinylpyridine was inactive as a genotoxic agent in the rat hepatocyte primary culture / DNA repair test when assayed at non-toxic doses.

Conclusions Remarks:

Conclusions of the authors (see References section).

Data Quality Reliability:

Klimisch Code = 2

Reliable with restrictions; basic data given: comparable to

guidelines/standards.

Data Quality Remarks:

not applicable

References:

Brunnemann, KD; Rivenson, A; Cheng, SC; Saa, V; Hoffmann, D. 1992. A study of tobacco carcinogenesis XLVII. Bioassays of vinylpyridines for genotoxicity and for tumorigenicity in A/J mice. *Cancer Letters*, 65, 107-

113.

Record Last Changed:

11/7/03

Order Number for Sorting:

34

General Remarks:

not applicable

Genetic Toxicity - In Vivo

ID: 30

Test Substance Identity: 2-Vinylpyridine

Test Substance Purity: > 99%

Test Substance Remarks: Test substance obtained from Aldrich Chemical and further distilled.

Method / Guideline Followed: not stated

Test Type: mouse lung adenoma assay

GLP: not stated

Year: 1992 Species: mouse

Strain: A/J strain, Jackson Laboratories, Bar Harbor, ME

Sex: female

Route of Administration: intraperitoneal injection

Doses / Concentration Levels: 200 µmol/mouse total dose

Exposure Period: injected 3 times weekly for total of 20 injections; 20 weeks post-dose

observation period

Statistical Methods: Student's t-test

Test Conditions Remarks: 4-(Methylnitrosamino)-1-(3-pyridyl)-1-butanone (NNK) was used as a

positive control.

2-Vinylpyridine was injected in 0.1 mL olive oil, using 23 mice. Negative

control (olive oil only) was also run for comparison.

Effect on Mitotic Index or PCE

/ NCE Ratio:

not stated

Genotoxic Effects: No induction of significant numbers of lung adenomas, or of any other

tumors, at a total dose of 200 µmol/mouse.

NOAEL (NOEL) / LOAEL

(LOEL):

not stated

Statistical Results: Lung tumor / mouse = 0.09 ± 0.28 , compared to 0.04 ± 0.20 for negative

control.

Results Remarks: not applicable

Conclusions: 2-Vinylpyridine is regarded as non-genotoxic and non-tumorigenic in strain

A/J mice.

Conclusions Remarks: Conclusions of the authors (see References section).

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; acceptable, well-documented publication/study

report which meets basic scientific principles.

Data Quality Remarks: not applicable

References: Brunnemann, KD; Rivenson, A; Cheng, SC; Saa, V; Hoffmann, D. 1992. A

study of tobacco carcinogenesis XLVII. Bioassays of vinylpyridines for genotoxicity and for tumorigenicity in A/J mice. Cancer Letters, 65, 107-

113.

Record Last Changed: 11/7/03

Order Number for Sorting:

General Remarks: not applicable

34

Repeated Dose Toxicity

ID:

31

Test Substance Identity:

2-Vinylpyridine

Test Substance Purity:

98.95%

Test Substance Remarks:

not applicable

Method / Guideline Followed:

not stated

Test Type:

repeated oral exposure

GLP:

yes

Year:

1983

Species:

rat

Strain:

not stated

Route of Administration:

gavage (corn oil carrier)

Duration of Test:

17 days

Doses / Concentration Levels:

13 doses over 17 day period

Sex:

male and female

Exposure Period:

17 davs

Frequency of Treatment:

once daily

Control Group and Treatment:

0 mg/kg of test compound

Post Exposure Observation

Period:

none

Statistical Methods:

not stated

Test Conditions Remarks

5 rats/sex/dose group; doses were 500, 200, 80 and 0 mg/kg. No data

available on age of animals.

Hematology for 80 and 200 mg/kg groups included RBC, Hgb., Hct., RBC

Ind., platelets, WBC, Diff.

Clinical chemistry for 80 and 200 mg/kg groups included AST (GOT), ADT

(GPT), SDH, AP, creatinine, UN, glucose.

NOAEL (NOEL):

NOEL < 80 mg/kg (males and females)

LOAEL (LOEL):

not determined

Actual Dose Received:

500, 200, 80, 0 mg/kg

Toxic Response / Effects:

500 mg/kg: Clinical signs: weakness, depressed activity, tremors, convulsions, sialorrhea. Gross pathology: edema of glandular stomach (females only), pallor of spleen (both sexes), enlarged and dark liver (females only). No data on weight gain, hematology, clinical chemistry or

organ weights due to early deaths.

200 mg/kg: Clinical signs generally the same as 500 mg/kg group. Histopathology: hyperkeratosis, acanthosis, hemorrhage, acute inflammation, edema and focal necrosis of nonglandular gastric mucosa in both males and females. Slight increase in liver weight in both sexes. Hematology: slight increase in atypical lymphocytes in females; slight increase in polymorphonuclear leucocytes in males; slight decrease in number of lymphocytes in males; all other parameters normal.

Clinical chemistry normal except for slight increase in ALT (GPT) in females. Weight gain and feed intake normal in females, slightly

depressed (during first 4 days) in males.

80 mg/kg: Clinical signs: sialorrhea in both males and females. Histopathology: hyperkeratosis and acanthosis of non-glandular stomach mucosa in both males and females; edema of gastric mucosa also observed in males only. Slight increase in liver weights in males only.

observed in males only. Slight increase in liver weights in males only. Hematology: slight increase in polymorphonuclear leucocytes in males only. Clinical chemistry all normal except for slight increase in AP in males.

Weight gain and feed intake normal for both sexes.

Statistical Results: not stated

Results Remarks: 500 mg/kg dose killed all rats after 1-2 treatments. Because of early

deaths, no tissue samples were taken.

Conclusions: Site of toxic action was non-glandular portion of the stomach (contact

tissue) and possibly the liver and central nervous system. The no effect

level was less than 80 mg/kg.

Conclusions Remarks: Conclusions of the authors (see References section).

Data Quality Reliability: Klimisch Code = 2

Reliable with restrictions; basic data given: comparable to guidelines /

standards.

Data Quality Remarks: not applicable

References: Anonymous. 1983. Basic Toxicity of 2-Vinylpyridine. Eastman Kodak

Company, Corporate Health and Environment Laboratories. As submitted to USEPA under TSCA §8(e) Compliance Audit Program agreement by

Eastman Kodak Company, 1992, OTS#0546362.

Record Last Changed: 11/5/03

Order Number for Sorting: 17

General Remarks: not applicable

Repeated Dose Toxicity

ID: 32

Test Substance Identity: 2-Vinylpyridine

Test Substance Purity: 97.34%

Test Substance Remarks: Test substance obtained from Reilly Tar & Chemical, lot number 30912.

When diluted in corn oil to concentrations of 0.8 to 7.2% and refrigerated,

mixture was stable for 11 days.

Method / Guideline Followed: Essentially the same as OPPTS 870.3100, "90-Day Oral Toxicity in

Rodents"

Test Type: 90-day repeated dose

GLP: yes
Year: 1983
Species: rat

Strain: Sprague-Dawley

Route of Administration: oral (gavage)

Duration of Test: 92 days

Doses / Concentration Levels: 180, 60, 20 and 0 mg/kg/day (7.2, 2.4, 0.8 and 0% solutions in corn oil)

Sex: both males and females

Exposure Period: 90 days

Frequency of Treatment: once daily, excluding weekends

Control Group and Treatment: yes

Post Exposure Observation none

Period:

.-

Statistical Methods: one-way analysis of variance (anova), Bartlett's test, and Duncan's multiple

range test. F-tests were performed where Bartlett's test indicated

significant difference in variances.

Test Conditions Remarks: 30 male and 30 female rats were assigned to each experimental group. All

animals were about six weeks of age at the start of the study. 10 randomly selected rats of each sex per dose were necropsied approximately halfway

through the study (43-day).

Ophthalmology exams were performed prior to star and during last week of

study.

Clinical pathology: Blood was collected at time of necropsy from the posterior vena cava under CO₂ anesthesia. Serum clinical chemistry tests

included:

Aspartate aminotransferase (AST)

• Alanine aminotransferase (ALT)

Alkaline phosphatase (AP)

Urea nitrogen

Glucose

Creatinine

Lactic dehydrogenase (LDH)

• γ-Glutamyl transpeptidase (GGTP)

Potassium

Calcium

Cholesterol

Sodium

Chloride

Phosphorus

Hematology tests included:

· Hemoglobin concentration

Hematocrit

Red blood cell count

· White blood cell count

· Differential white blood cell count

Platelet count

· Red blood cell indices

· Cell morphology

Necropsy: Rats were fasted overnight, anesthetized with CO₂ and exsanguinated by severing the posterior vena cava after collective blood samples. The following organs were weighed and their weights relative to body weight and to brain weight calculated:

• liver

kidneys

spleen

brain

heart

adrenal glands

ovaries

testes

The following organs were examined and collected in 10% buffered formalin. Eyes were fixed in a modified Zenker's (Russel's) fixative. All tissues from high and control groups were examined histologically. Target organs and gross lesions from other dose levels and interim kill groups were also examined histologically. (Examination of reproductive organs from this 90-day study meets the requirements for SIDS/HPV reproductive screening.)

trachea

lungs

aorta

tongue

esophagus

stomach

duodenum

jejunumileum

colon

coloncecum

rectum

urinary bladder

pituitary gland

pancreas

· thyroid gland

parathyroid glands

• thymus

· mesenteric lymph nodes

bone marrow

· cervical spinal cord

sciatic nerve

• eyes

skin

femur

• rib

salivary glands

vagina

uterus

• female mammary gland

testes

epididymides

· accessory sex organs (male)

male mammary gland

NOAEL (NOEL):

NOEL < 20 mg/kg/day for males; NOEL = 20 mg/kg/day for females

LOAEL (LOEL):

not stated

Actual Dose Received:

180, 60, 20 and 0 mg/kg/day

Toxic Response / Effects:

In male rats gavaged with 180 mg/kg/day, 2-vinylpyridine affected terminal body weight, absolute weight of liver, kidneys, brain, heart and adrenal glands, relative organ to body weight of liver, kidneys, brain, adrenal glands, and testes, and relative organ to brain weight of heart and adrenal glands.

In female rats gavaged with 180 mg/kg/day, 2-vinylpyridine affected terminal body weight, absolute weights of liver; and relative organ to body weights of liver, kidneys, and ovaries; and relative organ to brain weights of liver and ovaries.

Gavage with 60 mg/kg/day 2-vinylpyridine affected relative organ to body weights of liver and kidneys of male rats, relative liver to body and liver to brain weights of female rats.

Gavage with 20 mg/kg/day 2-vinylpyridine affected only the following weights in male rats: relative organ to body weights of kidneys and adrenal glands, absolute adrenal gland weight, and relative adrenal gland to brain weight.

After 92 days of experiment, gavage with 2-vinylpyridine, the principal effects were reduced body weights of high dose male rats to a greater degree than the female rats resulting in relative weight changes. No statistically significant effects on body weight gain of female rats was observed, but a slight reduction in weight gain of male rats given 180 mg/kg/day was seen, becoming statistically significant on day 4 of the study. The difference varied from 7.4% on day 21 to 13.8% on day 91. Male rats given 20 or 60 mg/kg/day had weight gains comparable to controls. In the female high dose group, weight reduction never exceeded 7%.

With the exception of day 57 in the 92-day group, high dose group males consistently ate less feed than controls, and the change was statistically significant. Lower dose group males (20 and 60 mg/kg/day) ate essentially identical amounts of feed as controls throughout the study, except for day 21 when the 20 mg/kg/day dose group ate less than the control group. On day 53 and 91, the 92-day female high dose group ate significantly more feed than controls, with differences of 14.2 and 11.1% respectively. No other statistically significant differences were observed in feed consumption.

Microscopically, the only compound-related effects observed were due to irritation of the gastric mucosa. These involved primarily the non-glandular epithelium and were characterized by degeneration of the superficial epithelial cells at the highest dose, hyperkeratosis and acanthosis of the epithelium resulting in a thickening of the non-glandular epithelium, and mild inflammatory changes (congestion, edema, and inflammatory cell infiltrates). Irritation of the glandular mucosa was generally much milder and not a consistent or dose-related effect.

Statistical Results: All reported results were analyzed for statistical significance.

Results Remarks: Only observations found to be statistically different from control groups are reported under "Toxic Response / Effects". All other examinations were

comparable to control groups.

Conclusions: The high dose (180 mg/kg/day) results in reduced body weight gain in

males, reduced feed consumption in males and also toward the end of the study in female rats; a slight increase in the number of platelets in both sexes, and a slight decrease in aspartate aminotransferase in male rats. This dose also affected male absolute weights of brain, heart, and adrenal glands, relative organ to body weight ratio of liver, kidney, brain, adrenal glands, and testes, and relative organ to brain weight ratio of heart and adrenal gland. In female rats, gavage with 180 mg/kg/day 2-vinylpyridine affected the absolute weight of the liver, and relative organ to body weight ratios of liver, kidneys and ovaries, and relative organ to brain weight ratios of liver and ovaries. The high dose (180 mg/kg/day) was clearly irritating to the non-glandular stomach epithelium of both sexes and microscopically was characterized by degeneration, hyperkeratosis and acanthosis resulting in a thickening of the non-glandular epithelium.

Conclusions Remarks: Conclusions of the authors (see References section).

Data Quality Reliability: Klimisch Code = 1

Reliable without restriction; comparable to guideline study.

Data Quality Remarks: not applicable

References: Vlaovic, Milan S. 1984. Subchronic Oral Toxicology of 2-Vinylpyridine.

Eastman Kodak Company, Toxicological Sciences Section, Health and Environment Laboratories. As submitted to USEPA under TSCA §8(e) Compliance Audit Program agreement by Eastman Kodak Company, 1992,

OTS#0546362.

Record Last Changed:

11/7/03

Order Number for Sorting:

17

General Remarks:

not applicable

Repeated Dose Toxicity

ID: 33

Test Substance Identity:

2-Vinylpyridine

Test Substance Purity:

98.3%

Test Substance Remarks:

Test substance obtained from Organic Synthetic Pharmaceutical Industries, lot 95-022. Stability of administration solution was checked at one week of storage in a refrigerator, showing no stability problems.

Method / Guideline Followed:

"Guidelines for 28-day Repeat Dose Toxicity Testing of Chemicals", Japan

Test Type:

28-day repeat dose

GLP:

yes

Year:

> 1991

Species:

rat

Strain:

Crj: CD (SD)

Route of Administration:

oral (gavage)

Duration of Test:

28-day exposure, up to 14-day recovery period

Doses / Concentration Levels:

0, 12.5, 50, 200 mg/kg/day

Sex:

males and females, 5 / sex / group

Exposure Period:

28 days

Frequency of Treatment:

once daily

Control Group and Treatment:

5 males; 5 females. Only corn oil was administered to the control group.

Post Exposure Observation

Period:

14-day recovery period for 0 and 200 mg/kg groups

Statistical Methods:

multiple comparison testing and Fisher probability computation

Test Conditions Remarks:

Recovery groups were employed for the 0 and 200 mg/kg dosages. Test substance was dissolved in corn oil and administered via gavage at about 0.5 mL per 100 g body weight. Animals were 6 weeks of age at study initiation. Body weight was measure once weekly through the end of the recovery period, and feed consumption was calculated weekly.

Clinical examination: Implemented twice (once at the end of administration and once at the end of recovery period).

Hematological analysis included:

white blood cell count

platelet number

red blood cell count

· white blood cell percentage

harrandable lavel

reticulocyte count

hemoglobin levelhematocrit level

prothrombin time

mean red blood cell volume

· activated partial thromblastin time

• mean red blood cell voidine

fibrinogen level

mean red blood cell hematochrome

mean red blood cell hematochrome concentration

Hematobiochemical analysis included:

total proteinalbumin

A/G ratio
 inorganic phosphorus

total bilirubin

calcium

blood sugar
 neutral fats
 total cholesterol
 urea nitrogen
 sodium
 potassium
 chlorine
 creatinine

• glutamic acid oxazaloacetate transaminase

glutamic acid pyruvic acid transaminase

· gamma-glutamyl transpeptidase

alkali phosphatase

Urinalysis included:

quantity
 color
 turbidity
 specific gravity
 sedimentation residue microscopy
 pH
 occult blood
 ketones
 sugar
 protein
 bilirubin
 urobilinogen

Necropsy: Animals were deprived of food for about 16 hours, anesthetized with ether, cut open at the abdomen and blood was collected from the large abdominal artery. Pathological examination included weights of the following organs:

brain
liver
kidneys
spleen
adrenals
testes
ovaries
thymus

Histopathology examination of control group and 200 mg/kg group included:

stomach
duodenum
kidneys
thymus
adrenals
heart
bone marrow (femur)

liver

Stomachs of medium and low dose groups were also examined.

NOAEL (NOEL):

NOEL = 12.5 mg/kg/day

LOAEL (LOEL):

not stated

Actual Dose Received:

0, 12.5, 50, 200 mg/kg/day

Toxic Response / Effects:

General Condition: Salivation was observed in both sexes receiving 50 and 200 mg/kg. All symptoms disappeared within hours of test substance administration.

Body Weight: Body weight gain was suppressed and food consumption decreased in males receiving 200 mg/kg. Once in the recovery period, a clear recovery trend was observed, and the increase in body weight was greater than in controls. Total feed consumption returned to control levels as of week 4 of administration.

Hematological examination: No difference was noted in either males or females in the examination of the control group and the groups administered the test substance. In males, the 200 mg/kg group had a

higher reticulocyte count than controls, but was still within the normal range of background values. In females, the 200 mg/kg group had a lower MCHC level than controls, but this was not a significant change, as there was no difference with controls in the MCV and MCH values computed.

Blood coagulation ability examination: At the ends of both the administration and recovery periods, no difference was found between the control group and the groups administered the test substance in either males or females.

Blood biochemical examination: At the end of the administration period, minor differences in certain biochemical levels were reported, sometimes in only one animal, but none were found to be statistically significant changes. No differences in any biochemical levels were found in either controls or administration groups in either males or females at the end of the recovery period.

Clinical Examination: Urinalysis at the end of administration revealed decreases in specific gravity in females receiving 50 and 200 mg/kg and volume increase in females receiving 200 mg/kg; these differences were not observed after the recovery period. No changes were observed in males vs. controls at any time.

Organ Weights and Ratios: Relative testes weights were increased in males receiving 200 mg/kg. Absolute and relative spleen weights were decreased and relative liver weights increased in females receiving 200 mg/kg. In both males and females, no differences in organ weights or ratios were observed at the end of the recovery period.

Pathological examination: Squamous hyperplasia and submucosal edema in the forestomach were observed in both sexes receiving 50 and 200 mg/kg, along with thickening of the mucosa at the higher dose. Moreover, erosion and cellular infiltration in the forestomach were observed in males receiving 200 mg/kg. Submucosal edema and/or erosion in the glandular stomach were also observed in females receiving 50 or 200 mg/kg. At the end of the recovery period, light squamous hyperplasia was observed in the forestomachs of three males and two females of the 200 mg/kg group.

Squamous hyperplasia in the forestomach was still observed in both sexes receiving 200 mg/kg at the end of the recovery period, but its incidence and extent were decreased in comparison to those of the same groups at the end of the administration period.

Statistical Results: See "Toxic Response / Effects"

Results Remarks: Only observations found to be statistically different from control groups are reported under "Toxic Response / Effects". All other examinations were

comparable to control groups.

Conclusions: Salivation was observed in both sexes receiving 50 and 200 mg/kg. Body weight gain was suppressed and food consumption decreased in males

receiving 200 mg/kg. Urinalysis revealed decreases in specific gravity in females receiving 50 and 200 mg/kg and volume increase in females receiving 200 mg/kg. Relative testes weights were increased in males receiving 200 mg/kg. Absolute and relative spleen weights were decreased and relative liver weights increased in females receiving 200 mg/kg. Squamous hyperplasia and submucosal edema in the forestomach were observed in both sexes receiving 50 and 200 mg/kg, along with thickening of the mucosa at the higher dose. Moreover, erosion and cellular infiltration in the forestomach were observed in males receiving 200 mg/kg. Submucosal edema and/or erosion in the glandular stomach were also observed in females receiving 50 or 200 mg/kg. Squamous hyperplasia in the forestomach was still observed in both sexes receiving 200 mg/kg at the end of the recovery period, but its incidence and extent were decreased in comparison to those of the same groups at the end of the administration period. The NOEL for repeat dose toxicity is considered

Conclusions Remarks: Conclusions of the authors (see References section).

to be 12.5 mg/kg/day for both sexes.

Data Quality Reliability: Klimisch Code = 1

Reliable without restriction; guideline study.

Data Quality Remarks:

not applicable

References:

Oba, Kousuke, et al. Twenty-eight day repeat dose oral toxicity test of 2-vinylpyridine in rats. Biosafety Research Center, Foods, Drugs and Pesticides (An-pyo Center), Japan, 582-2 Shioshinden Arahama, Fukudecho, Iwata-gun, Shizuoka, 437-12, Japan. (Located at Japan's Global

Information Network on Chemicals, found at http://wwwdb.mhlw.go.jp/ginc/index.html.)

Record Last Changed:

11/7/03

Order Number for Sorting:

54

General Remarks:

not applicable

Reproductive Toxicity

ID: 34

Test Substance Identity: 2-Vinylpyridine

Test Substance Purity: 97.34%

Test Substance Remarks: Test substance obtained from Reilly Tar & Chemical, lot number 30912.

When diluted in corn oil to concentrations of 0.8 to 7.2% and refrigerated,

mixture was stable for 11 days.

Method / Guideline Followed: essentially the same as OPPTS 870.3100, 90-Day Oral Toxicity in Rodents

Test Type: 90-day repeated dose

GLP: yes Year: 1983

Species: rat

Strain: Sprague-Dawley

Route of Administration: oral (gavage)

Duration of Test: 92 days

Doses / Concentration Levels: 180, 60, 20 and 0 mg/kg/day (7.2, 2.4, 0.8 and 0% solutions in corn oil)

Sex: both males and females

Duration of Test: 92 days

Frequency of Treatment: once daily, excluding weekends

Control Group and Treatment: see "Doses/Concentration Levels" above

Premating Exposure Period

(males):

not stated

Premating Exposure Period

(females):

not stated

Statistical Methods: one-way analysis of variance (anova), Bartlett's test, and Duncan's multiple

range test. F-tests were performed where Bartlett's test indicated

significant difference in variances.

Test Conditions Remarks: See Robust Summary #32 for test conditions details.

All tissues from high and control groups, including testes, epididymides, accessory male sex organs, ovaries, uterus, vagina, fallopian tubes and mammary glands (both sexes), were examined histologically. Target organs and gross lesions from other dose levels and interim kill groups

were also examined histologically.

NOAEL (NOEL): not stated

Actual Dose Received: 180, 60, 20 and 0 mg/kg/day

Toxic Response / Effects: Males:

Organ weights: There was a dose dependent increase in relative testes to body weights, however, only the high dose group reached statistical significance with a 20.3% increase. Statistical significance was also reached in relative testes to body weights in the 20 and 180 mg/kg/day dose groups killed mid-way through the experiment (at day 43), with 9.7%

and 14.6% increases, respectively.

Gross pathology: Enlarged prostate was observed in control group as well as some dose groups at day-92 (4/20 in control group; 3/20 in 20 mg/kg

group; 1/20 in 60 mg/kg group; 2/20 in 180 mg/kg group). No effects in epididymides or testes for any dose group.

Histopathology: No testicular effects observed any dose group. Chronic focal inflammation of the epididymides was observed in 4/20 rats in control group; 7/20 rats in 180 mg/kg group. Chronic inflammation of the prostate was observed in 6/20 rats in control group; 1/3 rats in 20 mg/kg group; and 9/20 rats in 180 mg/kg group.

Females:

Organ weights: In the 92-day rats, there was a statistically significant increase in relative ovary to body weight and ovary to brain weight in the high dose group with 24.1% and 18.2% increases, respectively.

Gross pathology: No effects at any dose level on fallopian tubes, vagina, uterus, mammary gland and ovaries. A single rat (0 mg/kg group at day 43) showed uterine hydrometra.

Histopathology: A single rat (1/20) at the 180 mg/kg dose level exhibited ovarian congestion; no other ovarian effects were observed at any dose level. No effects at any dose level were observed on fallopian tubes, vagina and female mammary gland. A single control group rat at 43-days showed uterine hydrometra.

Statistical Results:

All reported results were analyzed for statistical significance.

Results Remarks:

Additional results not related to reproductive organs are reported in Robust

Summary #32.

Conclusions:

A dose-dependent increase in relative testes to body weights was observed, but gross pathology and histopathology showed no testicular offects.

effects.

Likewise, a statistically significant increase in relative ovary weights was observed in the highest dose group, but again, no effects were observed upon gross pathology and histopathological examination.

No other pathology findings differed significantly from control groups, in either males or females.

Conclusions Remarks:

Conclusions of the authors (see References)

LOAEL (LOEL):

not stated

Data Quality Reliability:

Klimisch Code = 1

Reliable without restriction; comparable to guideline study.

Data Quality Remarks:

not applicable

References:

Vlaovic, Milan S. 1984. Subchronic Oral Toxicology of 2-Vinylpyridine. Eastman Kodak Company, Toxicological Sciences Section, Health and Environment Laboratories. As submitted to USEPA under TSCA §8(e) Compliance Audit Program agreement by Eastman Kodak Company, 1992,

OTS#0546362.

Record Last Changed:

12/4/03

Order Number for Sorting:

17R

General Remarks:

See Robust Summary #32 for additional testing details.

Per EPA guidance, an existing, adequate 90-day repeat dose study that "demonstrates no effects on reproductive organs, in particular the testes, then a developmental study (e.g., OECD Test Guideline 414) can be considered as an adequate test for information on reproduction / developmental effect." Thus, this 90-day study satisfies the requirements

for SIDS/HPV reproductive screening.